

Role of the Putative Polysaccharide Deacetylase BA1836 from *B. anthracis* in Spore Development and Germination

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How to cite this paper: Tomatsidou, A., Koutsoulis, D., Tzamarias, D., Kokkinidis, M., Vollmer, W. and Bouriotis, V. (2019) Role of the Putative Polysaccharide Deacetylase BA1836 from *B. anthracis* in Spore Development and Germination. *Advances in Microbiology*, 9, 679-702.

<https://doi.org/10.4236/aim.2019.98042>

Received: May 18, 2019

Accepted: August 4, 2019

Published: August 7, 2019

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Abstract

Putative and known polysaccharide deacetylases (PDAs) from *B. anthracis* have key roles in resistance to host lysozyme, stabilization of the cell wall, biogenesis of peptidoglycan (PG) and for neutral polysaccharide modification and attachment to PG. Here we elucidated the physiological role of the putative PDA BA1836 from *B. anthracis*. The *ba1836* gene was expressed upon entrance into the stationary phase of growth and enhanced during the early stages of sporulation. The $\Delta ba1836$ knockout strain had normal growth rate, did not exhibit any significant alterations in PG pattern of stationary phase cells and was not sensitive to lysozyme, but showed a defect in cell separation. Strikingly, the $\Delta ba1836$ mutant strain exhibited a severe delay in spore development although mature spores were ultimately developed and had normal morphology. Additionally, digestion of $\Delta ba1836$ mutant spore PG with mutanolysin produced an almost identical muropeptide pattern compared to peptidoglycan from wild type spores, although the amount of all muropeptides was significantly reduced. Finally, knockout spores exhibited a lower germination rate. To our knowledge, BA1836 has a unique role, among the presently characterized PDAs from *B. anthracis*, in spore development and germination.

Keywords

Bacillus anthracis, Polysaccharide Deacetylases, BA1836, Spores, Germination, Spore Development

1. Introduction

The Gram-positive bacterium *B. anthracis* exists in two distinct cellular morphologies: the vegetative cell and the dormant spore, which is the infectious agent of the anthrax disease [1]. Sporulation is a tightly regulated developmental process during which several morphological events take place [2].

Like in other *Bacillus* species sporulation in *B. anthracis* is induced by nutrient depletion of vegetative cells in combination with other stress signals [3] [4]. Initially, a single rod shaped cell divides asymmetrically, resulting in two different daughter cells, the mother cell and the forespore. Subsequently, the mother cell engulfs the forespore, in a process similar to phagocytosis, followed by the assembly of the spore coat and cortex. When the spore is mature, it is released into the environment, upon lysis of the mother cell, where the dormant spore is highly resistant to a range of environmental insults [2] [5] [6].

In vegetative cells, the net-like peptidoglycan (PG) layer is made of glycan chains connected by short peptides and is essential for cell integrity, enabling the cell to resist the turgor and maintain its cell shape [7]. Spore PG consists of two layers: the thin, inner germ cell wall and the thick outer cortex layer [8]. The cortex contains unique structural modifications, such as muramic-delta-lactam residues [9], and contributes to the resistance of the spores against heat and chemicals [10]. During germination, the cortex is specifically degraded by germination-specific lytic enzymes (GSLEs), while the germ cell wall remains intact. PG from spores lacking muramic-delta-lactam cannot be hydrolysed by GSLEs [11]. Enzymes and precursors employed for the production of the germ cell wall and cortex PG layers exhibit different localization. Those required for germ cell wall synthesis are localized in the forespore, while those involved in cortex synthesis are localized in the mother cell [12].

PG *N*-acetylglucosamine (GlcNAc) deacetylation has been demonstrated to contribute to lysozyme resistance of vegetative cells in several pathogenic species [13] [14] [15]. Deacetylation of spore PG has been also demonstrated, albeit at reduced levels compared to vegetative cells [16], but it is not associated with lysozyme resistance, since the main mechanism of spore lysozyme resistance is the permeability barrier of the spore coat layers [17].

After entry to a suitable host, the *B. anthracis* spore undergoes germination, which is the earliest pathogenic event, followed by the production of deadly toxins [18] [19]. Initiation of germination depends on interactions of small molecule nutrients, termed germinants, with receptors located within the inner endospore membrane [20]. Subsequently, calcium-dipicolinic acid (Ca-DPA) is released from the spore core thus initiating rehydration. Finally, GSLEs hydrolyse the spore cortex PG, leading to full rehydration of the core and resumption of metabolism [21] [22].

Polysaccharide deacetylases (PDAs) are members of the carbohydrate esterase family 4 (CE4) which consists of acetylxyylan esterases, chitin, chitooligosaccharide and PG deacetylases. Members of this family catalyze the hydrolysis of either *N*-linked acetyl group from GlcNAc residues (chitin, chitooligosaccha-

ride, and PG GlcNAc deacetylase) [13] [23] or *O*-linked acetyl group from *O*-acetylxylose residues (acetylxyloxy esterase) [24] [25] [26]. They are metal dependent enzymes and share a conserved catalytic core termed the NodB homology domain.

Interestingly, the genomes of *Bacillus* sp., and especially of *B. cereus sensu lato*, including *B. anthracis*, contain multiple putative PDA genes with high sequence homologies. The physiological role of seven of these from *B. anthracis* has been elucidated [27] [28]. BA1977 similarly to PG GlcNAc deacetylases (PGDAs) of other bacteria is involved in resistance to host lysozyme and required for full virulence. BA1961 and BA3679 participate in the biogenesis of the PG during both elongation and cell division, while BA5436 and BA2944 are important for neutral polysaccharide attachment to PG and for polysaccharide modification respectively [27]. Finally, BA0330 is important for the adaptation of the bacterium to grow in the presence of a high concentration of salt, whereas BA0331 contributes to the maintenance of a uniform cell shape [28]. Due to their crucial roles in *B. anthracis* physiology, inhibitors targeting their conserved active sites have been developed [29].

Presently, there is limited information on PDAs affecting either sporulation or germination of bacteria. Two PDAs from *B. subtilis*, PdaA and PdaB have been demonstrated to affect sporulation and germination of this bacterium. Absence of PdaA, a PG *N*-acetylmuramic acid (MurNAc) deacetylase from *B. subtilis*, results in complete lack of spore germination due to deficiency in muramic- δ -lactam formation, one of the major components of the spore cortex [30]. The absence of PdaB results in empty or cortex-less spores, implying its importance in cortex formation and further maintenance of spores after the late stage of sporulation [31].

In this study we used genetic (knock-out) analysis, protein localization as well as spore and germination assays to elucidate the function of the putative PDA BA1836 from *B. anthracis*. We show that this protein has a unique role among the presently characterized putative and known PDAs of *B. anthracis* in promoting normal spore development and germination.

2. Materials and Methods

2.1. Bacterial Strains and Growth Conditions

The strains and plasmids used in this study are listed in **Table S1**. *E. coli* strains were grown in Luria Bertani (LB) medium and *B. anthracis* strains in Brain Heart Infusion (BHI) medium. Sporulation was induced by resuspension in modified G medium [32] [33] and mature spores were formed after plating of vegetative cells on Spizizen Yeast (SPY) agar medium which when required was supplemented with 0.2% xylose.

2.2. Construction of *B. anthracis* Δ ba1836 Knockout Strain, GFP Fusion and *lacZ* Transcriptional Fusion

A strain bearing an in-frame deletion of *ba1836* gene was created using a mar-

kerless gene replacement method [34]. In brief, DNA fragments containing the sequence upstream and downstream of *ba1836* gene were amplified from genomic DNA of *B. anthracis* 7702 [35] by PCR using the appropriate oligonucleotides (Table S2). Both fragments were cloned into the same pGEM vector and the upstream-downstream fragment was digested with SacI/KpnI and ligated into pRP1028 vector. *E. coli* HB101 [36] was used as helper strain in order to primarily introduce pRP1028 construct [37] and subsequently pSS4332 vector [37], which encodes the I-SceI enzyme, into *B. anthracis* 7702 strain. Finally, the knockout strain had no resistance to antibiotics and was validated for the correct construction by PCR.

To construct strains expressing GFP translational fusions, the *gfp-mut1* gene was amplified from pNF8 [38] with specific primers, digested with NruI and BglII, and ligated into the xylose-inducible plasmid pWH1520 [39]. Then, *ba1836* (lacking the stop codon) was amplified from *B. anthracis* 7702 chromosomal DNA with the appropriate primers to incorporate at the C terminus the polylinker GPGP. The amplicon was digested with BamHI and NruI and ligated in-frame to the 5' end of *gfp-mut1*, placing the gene fusion under a xylose-inducible promoter. *E. coli* GM48 (*dam*[−]) was transformed with the resulting construct to obtain nonmethylated plasmid DNA for electroporation [40] into 7702 and Δ *ba1836* *B. anthracis* cells. 10 ml of BHI medium were inoculated from overnight cultures to an OD_{600 nm} of 0.05 and incubated at 37°C. For *ba1836-gfp* expressing strains, induction was achieved with 0.2% xylose (for expression from the xylose-inducible promoter) 1 h before the culture reached early stationary phase of growth (OD_{600 nm} around 3.0).

In order to create the *lacZ*-transcriptional fusion, a DNA fragment of the upstream region of *ba1836* gene was amplified from *B. anthracis* 7702 chromosomal DNA. Primers were synthesized to incorporate a HindIII site at the start and a BamHI site at the end of the amplified fragment (Table S2), which was digested with the corresponding enzymes, and ligated into the multicopy replicative vector pHT304-18Z [41]. In this way, a *lacZ* reporter construct with the upstream region of each gene fused directly to the start codon of the *lacZ* gene was obtained. *E. coli* GM48 (*dam*[−]) was transformed with the resulting construct to obtain nonmethylated plasmid DNA for electroporation [40] into 7702 *B. anthracis* cells.

2.3. Site-Directed Mutagenesis of *ba1836* Gene

A mutant of the *ba1836* gene was constructed using a two-step/four-primer overlap extension PCR method [42]. Primers used in this method are listed in Table S2. The amplified *ba1836* product was cloned into pWH1520 vector as described above.

2.4. Preparation of Sporulating Cell Cultures

10 ml of BHI medium were inoculated from overnight cultures of 7702 parental

and $\Delta ba1836$ mutant strains to an $OD_{600\text{ nm}}$ of 0.05 and incubated at 37°C. When $OD_{600\text{ nm}}$ of the cultures reached 1.2 (mid-exponential phase) or 3.0 (early stationary phase), the cultures were centrifuged for 20 min at 2570 g and sporulation was induced by resuspension of the cell pellet in an equal volume of modified G medium. The cultures were then transferred to 30°C. Cells were collected at frequent time intervals and observed under optical and fluorescence microscopy.

2.5. β -Galactosidase Activity Assay

Cell samples were collected in triplicates from shaking cultures at various time points during vegetative growth or sporulation. $OD_{600\text{ nm}}$ for each cell sample was recorded before collection. β -galactosidase activity was measured as described by Camp and Losick [43] with minor changes. Cells pellets corresponding to 1 ml of the original sporulation medium were resuspended in 0.2 ml Z-buffer (60 mM Na_2HPO_4 , 40 mM NaH_2PO_4 , 10 mM KCl, 1 mM $MgSO_4$, 50 mM β -mercaptoethanol pH 7.0). 50 μ l of the cell suspension were added to individual wells of a 96-well plate containing 15 μ l of 0.1% SDS and were incubated at 37°C for 30 min. 20 μ l of 4 $mg\cdot mL^{-1}$ 2-nitrophenyl β -D-galactopyranoside (ONPG) in Z-buffer were then added to each well and mixed thoroughly. Absorbance at 420 nm for each reaction was read once per minute for 1 h at 37°C in a Fluostar Galaxy plate reader. β -galactosidase activity is expressed in Miller units (MU) [44] [45].

2.6. Fluorescence Microscopy, Scanning and Transmission Electron Microscopy

B. anthracis sporulating cell cultures of 7702 parental and $\Delta ba1836$ knockout strains were examined by optical and fluorescence microscopy. For membrane visualization the FM4-64 fluorescent dye was added to a final concentration of 1 $\mu g\cdot mL^{-1}$. Sporulating cells in stages II and III were stained with FM4-64 in order to visualize the asymmetric septum and engulfment respectively. Optical microscopy was used after stage III of sporulation when the forespore is formed. *B. anthracis* vegetative cell cultures of 7702 overexpressing the *ba1836-gfp* fusion were examined by fluorescence microscopy. The images were obtained without fixation on an inverted epifluorescence microscope Nikon E800. Preparation of cell samples for scanning and transmission electron microscopy was performed according to Arnaouteli *et al.* [28]. Samples were examined using a JEOL JSM-6390LV scanning electron microscope, operating at 20kV. Sections were observed in a JEOL, JEM 2100 transmission electron microscope, operated at 80 kV.

2.7. In Vitro Determination of Lysozyme Sensitivity

To test the sensitivity of $\Delta ba1836$ mutant strain in the presence of exogenously added lysozyme, overnight cultures of the parental and mutant strains were diluted to an $OD_{600\text{ nm}}$ of 0.1 into 2 cultures of 20 ml of fresh SPY medium for each strain. The cultures were incubated at 37°C until an $OD_{600\text{ nm}}$ of 1.2 and 3.0 cor-

responding to mid-exponential and early stationary phase of growth respectively. Then each culture was divided in two equal parts of 10 ml, and 10 $\mu\text{g}\cdot\text{ml}^{-1}$ hen egg lysozyme was added to one of the two subcultures. The growth of both treated and untreated subcultures was monitored.

2.8. Autolysis Assay

Autolysis assay was performed according to Balomenou *et al.* [27] with minor changes. Cultures of 10 ml SPY of 7702 and $\Delta ba1836$ cells at an $\text{OD}_{600\text{ nm}}$ of 3.0 were centrifuged for 20 min at 2570 g and the bacterial pellet was concentrated 10 times in SPY medium to which 10 mM NaN_3 was added. The decrease of $\text{OD}_{600\text{ nm}}$ of each culture was monitored every 30 min for a period of 180 min.

2.9. PG Purification

PG from the parental strain 7702 and $\Delta ba1836$ mutant was purified from stationary growing bacteria (in SPY) as well as from spores, as previously described [46] [47]. Muropeptides were generated from 0.2 mg of spore or vegetative cell PG by digestion with mutanolysin (Sigma) at 37°C for 16 h and separated by HPLC as previously described [48].

2.10. Competition Experiments

In order to perform competition experiments, a *lacZ* reporter construct with the *eag* promoter (*Peag*) fused directly to the start codon of the *lacZ* gene was built. The *Peag* [49] was amplified from *B. anthracis* 7702 chromosomal DNA. Primers were synthesized to incorporate a HindIII site at the start and a BamHI site at the end of the *Peag* (Table S2), which was digested with the corresponding enzymes, and ligated into the multicopy, replicative vector pHT304-18Z. *E. coli* GM48 (*dam*[−]) was transformed with the resulting construct to obtain nonmethylated plasmid DNA for electroporation [40] into 7702 *B. anthracis* cells.

Competition experiments were performed according to Traag *et al.* [45] with a few changes. Briefly, starting from overnight BHI culture, 7702 strain containing pHT304-18Z-*Peag* construct was mixed, either in 1:1 or 1:3 ratio with mutant $\Delta ba1836$ strain containing pHT304-18Z vector in 5 ml modified G medium. Cultures were grown and allowed to sporulate at 30°C for 48 h. The cultures were then heat treated at 80°C for 20 min, briefly cooled at room temperature and inoculated in 5 ml fresh modified G medium. Before inoculation, cultures were diluted and plated on agar plates containing 40 $\mu\text{g}\cdot\text{ml}^{-1}$ 5-bromo-4-chloro-3-indolyl- β -D-galactopyranoside (X-Gal) and 5 $\mu\text{g}\cdot\text{ml}^{-1}$ erythromycin. The blue and white colonies were counted. Similarly, competition experiments were performed for vegetative phase of growth. Briefly, starting from overnight BHI culture, 7702 strain containing pHT304-18Z-*Peag* construct was mixed in 1:1 ratio with mutant $\Delta ba1836$ strain containing pHT304-18Z vector in 5 ml BHI medium. Cultures were grown at 30°C for 24 h and then inoculated in 5 ml fresh BHI medium [45].

2.11. Spore Preparation and Purification

Single colonies from 7702 and $\Delta ba1836$ *B. anthracis* strains were streaked onto SPY agar plates supplemented with 0.2% glucose and 10 mM $MgSO_4$ and left to sporulate for 7 days at 30°C. When needed, the SPY agar plates were additionally supplemented with 0.2% xylose.

Spores were harvested in water, centrifuged at 5900 g for 5 min, washed twice in water, heated at 65°C for 20 min, washed again and stored at 4°C. Microscopic examination indicated that all spore preparations were 98% free of contaminating vegetative cells, sporulating cells and germinated spores.

2.12. Spore Viability

Spore viability was defined as the number of colony-forming units (cfu) formed compared to the number of phase-bright spore particles plated. Purified parental 7702 and $\Delta ba1836$ spores were tittered by visual counting using a Neubauer haemocytometer (spores/ml), then diluted and plated onto LB agar, with or without prior heat treatment at 80°C for 20 min, to measure cfu (cfu/ml) [50].

2.13. Spore Germination

Spore germination was performed according to Fazzini *et al.* [51] with certain modifications. Briefly, spores were suspended at an $OD_{600\text{ nm}}$ of 1.0 in 1 ml water. After heat activation at 65°C for 20 min, the suspensions were centrifuged at 5900 g for 5 min and 900 μ l were removed. Germination of spores was initiated by the addition of 1ml of a mixture of 25 mM L-alanine and 12.5 mM inosine. The suspensions were incubated at 37°C for 120 min. Germination was monitored by the loss of optical density at 600 nm.

2.14. Western Blotting Analysis

B. anthracis cell lysates, at the time point at which GFP fluorescence signal was obtained, were separated by SDS-PAGE, blotted, and probed with the following antibodies: polyclonal rabbit anti-GFP primary antibody (Minotech) diluted 1:25,000 and monoclonal anti-rabbit IgG (γ -chain specific) alkaline phosphatase secondary antibody (Sigma) diluted 1:3125.

3. Results

3.1. BA1836 Is a Putative PDA

The genome of *B. anthracis* Sterne 7702 sequence database revealed 12 coding sequences for putative PDAs of the CE4 family. Eleven of them exhibit more than 90% sequence identity to their homologues from *B. cereus sensu stricto*. For BA1836, the TMHMM [52] program predicts the presence of an *N*-terminus transmembrane domain (residues 5 - 23). The TatP [7] [53] and SignalP [53] [54] programs do not predict a signal peptide. BA1836 shares 92% identity with its homologue BC1768 from *B. cereus* and displays 25% - 30% identity with *B. subtilis* PDAs. Additionally, BA1836 shares an approximately 30% identity with

other putative and known PDAs from *B. anthracis*.

Importantly, the sequence alignment of BA1836 (excluding residues 1 - 47) with representative members of the CE4 family [24] shows that BA1836 retains the pattern of sequence motifs (MT1-MT5) which are typical for PDAs (Figure 1). These include motif MT2, which is associated with a metal cation binding site formed in the catalytic center of the enzyme, and key catalytic residues (MT1, MT5), which occupy structurally homologous positions in CE4 esterases of known structures [55]. More specifically, Asp⁵³ of motif MT1 and His¹⁹⁶ of motif MT5, corresponding to the catalytic base and the catalytic acid of the deacetylation reaction respectively, are conserved in the sequence of BA1836 (Figure 1). Additionally, the highly conserved Pro¹⁴⁵, which we have recently demonstrated to be autocatalytically hydroxylated in the presence of molecular oxygen at the Ca atom to produce 2-hydroxyproline (2-Hyp), is found in motif MT3, in agreement with other well characterized PDAs. The self-hydroxylation of proline at the Ca atom has been shown to significantly enhance enzymatic activity [56]. The pattern of hydrophobic residues in motifs MT4 and MT5 agrees well with the patterns found in other PDAs (Figure 1).

3.2. BA1836 Is Expressed in Stationary Phase Cells and Further Induced during Sporulation

In order to monitor the expression of the *bal836* gene we constructed a transcriptional reporter by fusing an upstream fragment of *bal836* (containing the

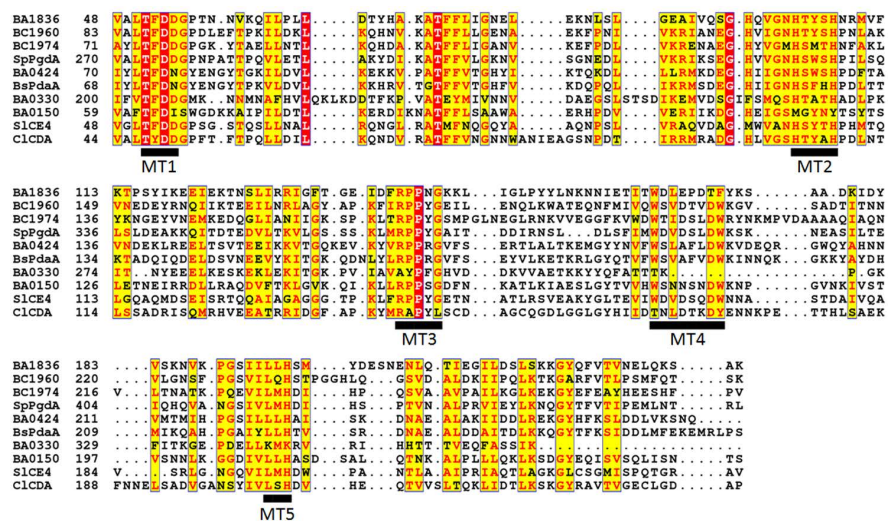


Figure 1. Sequence alignment of NodB domains. Sequence alignment of the NodB domain of BA1836 with representative members of the CE4 family including the PGDAs BC1960 [23] [57], BC1974 (*B. cereus*) [27] [58] and SpPgda (*S. pneumoniae*) [13] [55], the MurNac peptidoglycan deacetylases BA0424 (*B. anthracis*) [59] and BsPdaA (*B. subtilis*) [30] [60], the putative PDA BA0330 (*B. anthracis*) [28], the inactive PDA BA0150 (*B. anthracis*) [61], the acetylxylen esterase SICE4 (*S. lividans*) [62] and the chitin deacetylase ClCDA (*C. lindemuthianum*) [63]. The conserved sequence motifs MT1-MT5 for this class of enzymes are shown at the bottom. The alignment was performed with T-coffee and plotted with the ESPRIT [64].

promoter region) with the start codon of the *E. coli lacZ* gene. 7702 *B. anthracis* cells carrying this construct were harvested at various time points during vegetative growth. As shown in **Figure 2(a)**, BA1836 promoter driven *lacZ* expression was absent during exponential growth, but was rapidly induced to high level shortly after cells entered the stationary phase of growth.

We next resuspended early stationary phase 7702 *B. anthracis* cells carrying this construct into sporulation-inducing medium and collected samples at various time points during sporulation. As we observed by optical microscopy, most cells were in stage II of sporulation (asymmetric division) about 8 - 12 h after transfer into sporulation medium, in stage III (engulfment) after 12 - 16 hours, in stage IV-V (coat and cortex assembly) after 16 - 20 hours and in stage VI-VII (spore maturation and mother cell lysis) after 20 - 24 hours. We observed that expression of *ba1836-lacZ* further increases during the early stages of sporulation (Stage I-II) (**Figure 2(b)**). In contrast, when exponentially growing 7702 *B. anthracis* cells, in which *ba1836-lacZ* was not expressed, were induced to sporulate, the BA1836 promoter remained silent (**Figure S1**). This suggests that sporulation conditions alone are not sufficient to induce BA1836 expression.

3.3. BA1836-GFP Localizes to the Cell Periphery and Septa

The amino acid sequence of BA1836 is predicted to include an *N*-terminal membrane spanning region that is not removed by signal peptidase. In order to determine the subcellular localization of BA1836, a C-terminal *gfp* fusion of the *ba1836* gene was constructed and introduced into *B. anthracis* 7702. We induced the expression of *ba1836-gfp* in *B. anthracis* cells that entered the stationary phase of growth, *i.e.*, when BA1836 was normally expressed. Expression of the *gfp*-fused BA1836 was confirmed by Western blot analysis (**Figure S2**). The GFP tag did not affect the function of the protein, because the mutant strain transformed

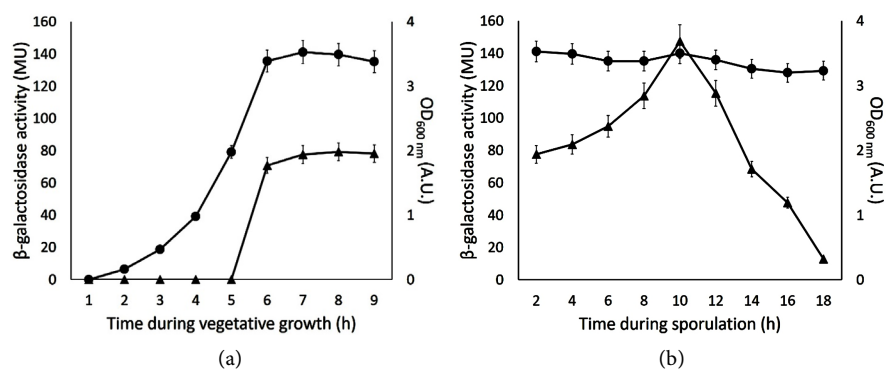


Figure 2. Expression of *ba1836* gene. (a) *B. anthracis* 7702 strain, carrying the *lacZ* transcriptional reporter for *ba1836* gene, grown in BHI (b) induction of sporulation of stationary phase cells. Samples taken at the indicated time points were assayed for β -galactosidase activity. *ba1836* is expressed after cells entered stationary phase of growth and is further induced in sporulation. (●) OD_{600 nm} (A.U.) (▲) β -galactosidase activity in Miller Units (MU). Data are representative of three independent experiments, each experiment was performed in triplicates and the error bars indicate standard deviation.

with the respective BA1836-GFP protein complemented the observed phenotypes (section: $\Delta ba1836$ cells display severe delay in spore development and $\Delta ba1836$ knockout strain lowers the spore germination rate). BA1836-GFP localized at the cell membrane and septa in stationary phase cells (**Figure 3**).

3.4. Cells Lacking BA1836 Suffer Reduced Sporulation Fitness

To elucidate the physiological role of BA1836, we constructed a *ba1836* gene knockout in *B. anthracis* 7702 strain according to the method of Janes and Stibitz [34]. Mutant cells were able to grow with no obvious defect and with similar rate as wild-type cells in various liquid media (BHI and SPY broth) indicating that BA1836 is not required for growth and viability of *B. anthracis* cells (**Figure S3(a)**, **Figure S3(b)**). $\Delta ba1836$ knockout cells grown in SPY medium, were not sensitive to lysozyme either in the exponential or in the stationary phase of growth (**Figure S4(a)**) suggesting that BA1836 is not a *bona fide* PGDA involved in resistance to host lysozyme. The autolytic activity of $\Delta ba1836$ and 7702 strains was tested by addition of NaN_3 , a known inducer of autolysis in growing cultures. $\Delta ba1836$ cells showed decreased autolysis at these conditions compared to the parental strain (**Figure S4(b)**). The PG pattern from stationary phase cells of $\Delta ba1836$ mutant was similar to that of the parental strain (**Figure S5**).

In order to examine the morphology of $\Delta ba1836$ cells, we collected samples during exponential and stationary growth phase in SPY medium and visualized cells by scanning electron microscopy (SEM). $\Delta ba1836$ daughter cells grown exponentially displayed morphologies similar to that of the parental strain, consistent with the lack of *ba1836* expression under these conditions (**Figure S4(c)**). However, stationary phase $\Delta ba1836$ cells displayed chains of unseparated cells ($5.15 \pm 0.14 \mu\text{m}$, $n = 190$) compared to the mostly single cell morphology ($3.21 \pm 0.18 \mu\text{m}$, $n = 175$) of the parental strain (**Figure S4(c)**).

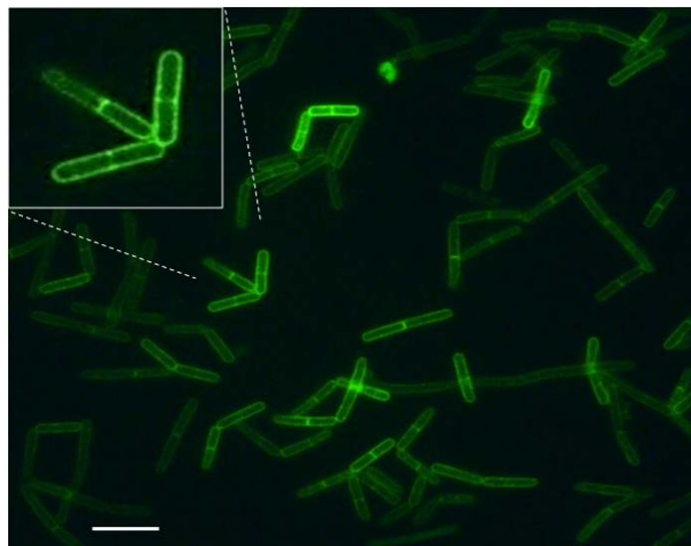


Figure 3. Localization of BA1836-GFP. BA1836-GFP localizes at the cell periphery and septa in stationary phase cells. Scale bar 10 μm .

Since *ba1836* gene was expressed upon entry to the stationary phase of growth and was further induced during sporulation, we next examined the morphology of spores produced by $\Delta ba1836$ by SEM and TEM imaging. No significant differences were observed between mature spores of $\Delta ba1836$ and the parental strain, suggesting that $\Delta ba1836$ cells can produce morphologically and structurally normal spores with similar cortex thickness (**Figure S6(a)**). The efficiency of $\Delta ba1836$ spore formation, their viability and their heat resistance was similar to that of the parental strain (**Figure S6(b)**). To test for a more subtle role of *ba1836* in spore formation, competition experiments were conducted in which $\Delta ba1836$ strain competed against 7702 wild type strain for several rounds of sporulation. In 7702 wild type strain, *lacZ* is under the transcriptional control of the *eag* promoter. The competition experiment was initially performed with a ratio of 1:1 (wild type to mutant cells) (**Figure 4(a)**) but since $\Delta ba1836$ cells showed a severe competition deficit, the mutant cells were added in threefold excess (**Figure 4(b)**). Over the course of seven days of co-culturing the percentage of the mutant strain decreased from 75% to 1% of the population (**Figure 4(b)**). On the contrary, $\Delta ba1836$ mutant strain had no fitness defect under vegetative growth conditions (**Figure 4(c)**). Complementation of the competition deficit under sporulation-inducing conditions of $\Delta ba1836$ strain could not be performed since the addition of xylose to the cultures inhibited entering of cells to sporulation. These results strongly suggest that BA1836 is actively involved in sporulation and/or germination processes.

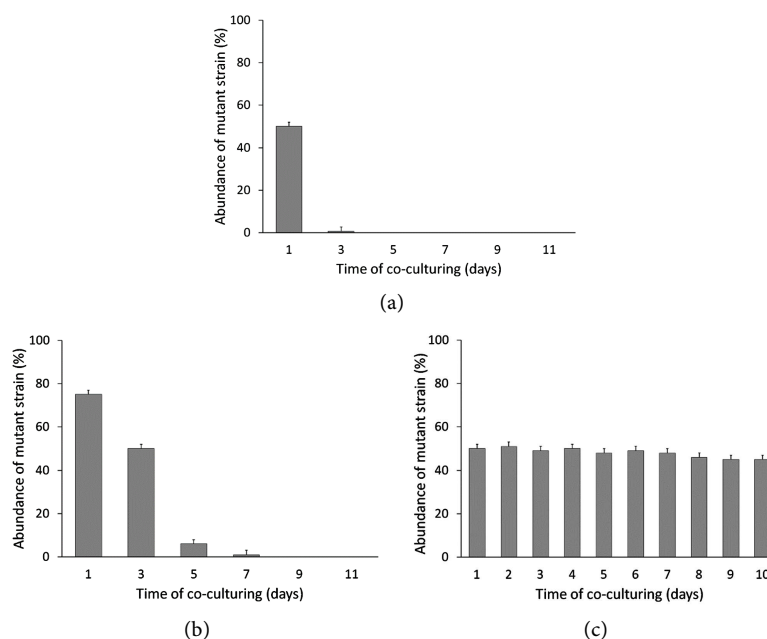


Figure 4. Competition experiment. $\Delta ba1836$ mutant strain exhibits reduced sporulation fitness. A wild-type reference strain carrying a *Peag-lacZ* transcriptional reporter was competed against $\Delta ba1836$ mutant in sporulation—inducing medium, starting with (a) 50% and (b) 75% mutant strain; (c) $\Delta ba1836$ mutant strain has no fitness defect under growth conditions. Data are representative of three independent experiments, each experiment was performed in triplicates and the error bars indicate standard deviation.

3.5. $\Delta ba1836$ Cells Display Severe Delay in Spore Development

Since we observed a competition deficit under sporulation—inducing conditions of $\Delta ba1836$ knockout cells, we next examined every stage of sporulation by optical microscopy. Interestingly, the mutant cells showed a remarkable delay in spore development (**Figure 5**). At 13 hours post-resuspension into sporulation—inducing medium, the majority of 7702 mother cells have engulfed the forespore (stage III of sporulation). In contrast, at the same time point the majority of the $\Delta ba1836$ cells only entered assymetric septation (stage II of sporulation) (**Figure 5**). Four hours later, almost every 7702 mother cell has produced a mature forespore (stage VI of sporulation), while the majority of the $\Delta ba1836$ mother cells have only entered stage III of sporulation (**Figure 5**). Mature forespores were visible in $\Delta ba1836$ mother cells 21 hours post resuspension into sporulation—inducing medium (**Figure 5**). These observations show that $\Delta ba1836$ cells display a severe delay in spore development, consistent with the expression of *ba1836* in this developmental stage.

The PG pattern from mature spores of $\Delta ba1836$ and 7702 strains were analyzed by HPLC [47]. The digestion of PG from $\Delta ba1836$ spores with mutanolysin produced an identical overall mucopeptide profile compared to PG from wild type spores, although the amount of mucopeptides released from the mutant was significantly reduced (**Figure 6**). Expressing *ba1836-gfp* in $\Delta ba1836$ restored the wild-type spore PG profile (**Figure 6**). Complementation was also attempted using the native protein but no expression was observed in $\Delta ba1836$ cells (data not shown).

3.6. $\Delta ba1836$ Knockout Strain Lowers the Spore Germination Rate

Finally, we determined the rate of germination of $\Delta ba1836$ mutant strain as a

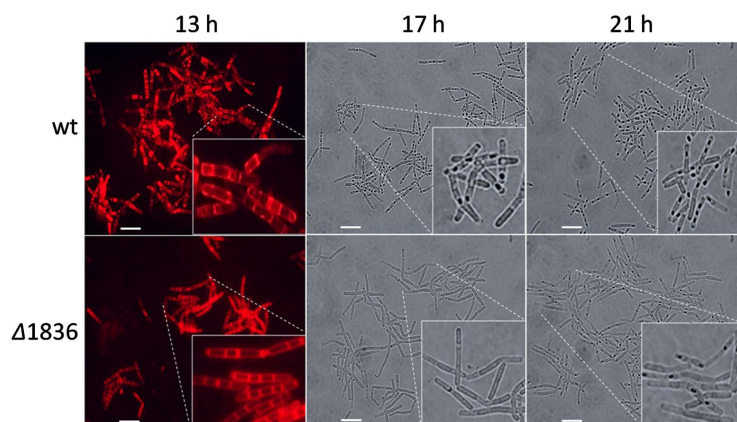


Figure 5. Phenotypic analysis of parental and mutant sporulating cells. Optical microscopy of 7702 and $\Delta ba1836$ cells, at various time points after induction of sporulation into G modified medium. According to our observation, 8 - 12 hours corresponded to Stage II, 12 - 16 hours to Stage III, 16 - 20 hours to Stage IV-V and 20 - 24 hours to Stage VI of sporulation. $\Delta ba1836$ shows a delay of spore development of approximately 4 hours. Scale bar 10 μ m. Data are representative of three independent experiments.

reduction in spore-specific absorbance upon the addition of 25 mM L-alanine and 12.5 mM inosine as germinants. As shown in **Figure 7**, when *ba1836* gene was deleted, spores were capable of initiating germination but, compared to wild-type, the $\Delta ba1836$ spores had a lower rate of germination. Fifteen minutes after the addition of germinants there was almost 40% loss of absorbance for 7702 spores compared to a 20% loss for $\Delta ba1836$ spores. $\Delta ba1836$ spores lost almost 40% of the absorbance after 90 min. These results indicate that the absence of BA1836 results in a delay in germination. The rate of germination of $\Delta ba1836$ mutant spores was partially complemented by expressing either BA1836-GFP or the predicted catalytically inactive BA1836 D53A-GFP, indicating that putative deacetylase activity is not required to complement this phenotype (**Figure 7**).

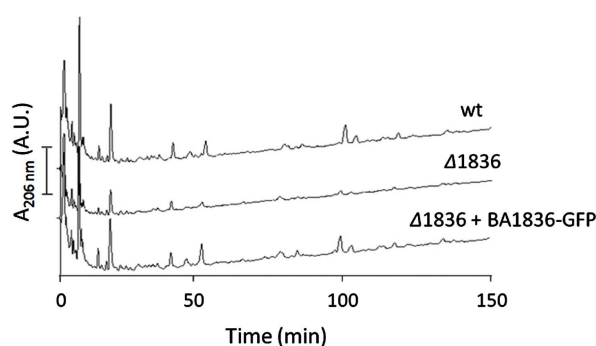


Figure 6. HPLC analysis of mucopeptide pattern of peptidoglycan isolated from 7702 and $\Delta ba1836$ mature spores. $\Delta ba1836$ spore PG digestion with the muramidase mutanolysin produced an identical overall mucopeptide profile but reduced amount of mucopeptides. $\Delta ba1836$ mutant strain complemented with BA1836-GFP recovers the phenotype. Bar, 200 mAU. Data are representative of three independent experiments.

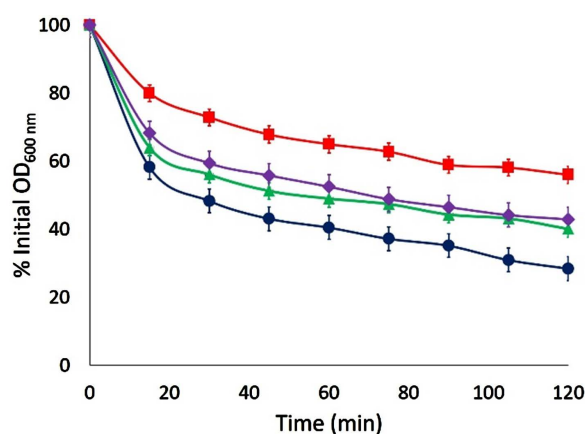


Figure 7. Germination of parental and mutant spores in response to L-ala and inosine. Spores of 7702 (●) and $\Delta ba1836$ (■) were induced to germinate by the addition of L-ala and inosine. Germination was monitored by loss of absorbance at 600 nm. $\Delta ba1836$ shows a decreased rate of germination compared to the wild type. Mutant strain complemented with BA1836-GFP (▲) and BA1836 D53A-GFP (◆) partially recovered the phenotype. Data are representative of three independent experiments, each experiment was performed in triplicates and the error bars indicate standard deviation.

4. Discussions

In this study we employed gene knock-out, protein localization, expression profile analysis as well as sporulation and germination assays to elucidate the physiological role of the putative PDA BA1836 from *B. anthracis*.

ba1836 expression is triggered upon entrance in the stationary phase of growth and is further induced in the early stages of sporulation, in contrast to a *B. anthracis* DNA microarray study which reported that *ba1836* is expressed only throughout logarithmic phase of growth [65].

The $\Delta ba1836$ mutant showed a defect in cell separation (**Figure S4(a)**) and a lower rate of autolysis (**Figure S4(b)**) compared to the wild type strain suggesting a mechanism of impaired substrate recognition by PG hydrolases that are crucial for autolysis and cell division. Reduced lysis has been similarly observed with all PDA mutants of *B. anthracis* examined so far, except $\Delta ba1977$ [27].

Deletion of *ba1836* resulted in a severe competition deficit under sporulation—inducing conditions (**Figure 4(a)** and **Figure 4(b)**). $\Delta ba1836$ cells displayed a severe delay in sporulation (**Figure 5**), although mutant spores were morphologically and structurally normal and able to germinate (**Figure S6(a)** and **Figure S6(b)**).

Deletion of other genes from *B. anthracis* has been similarly reported to result in a noticeable delay in spore development which has been attributed to different reasons. Inactivation of almost each of the sensor histidine kinase genes in *B. anthracis* resulted in a delay in sporulation of the resulting mutant strains, especially the $\Delta ba2291$ strain [66]. Similarly to *ba1836*, *ba2291* was expressed during stationary phase of growth and was inferred to be involved in sporulation by its strong inhibition of sporulation when expressed in *B. subtilis*. The inhibitory activity of BA2291 has been proposed to be due to either dephosphorylation of the phosphotransferase Spo0F or the transcription factor Spo0A or sequestering one or both of these proteins, preventing access by sporulation kinases [66]. Similarly one of the three sortase mutant strains of *B. anthracis*, namely *srtC* mutant, did not readily form spores in guinea pig tissue or sheep blood unless their vegetative forms were expressed in the presence of oxygen [67]. It is currently unclear if BA1836 is functionally linked to BA2291 or SrtC.

In repeated mutanolysin digests of spore PG from $\Delta ba1836$ mutant, we obtained an almost identical muropeptide pattern, but a considerable lower amount of muropeptides, compared to digests of spore PG from wild-type. This could be due to a reduced amount of PG in the mutant, less efficient digestions, or both (**Figure 6**). The wild-type spore PG profile was partially complemented when the mutant strain was transformed with BA1836-GFP. A similar phenotype has been reported for the sporulation proteins SpoVM and SpoVS of *B. subtilis* [68]. The reduced PG amount of *spoVM* deficient spores has been attributed to incomplete cortex synthesis which was suggested to result either from its interaction with the spore coat morphogenesis protein SpoIVA [69] or the protease FtsH [70] [71]. Except from the reduced amount of spore PG, *spoVS*—deficient

spores showed a delay in the initiation of sporulation. Similarly to SpoVM and SpoVS, BA1836 might have a regulatory role via its interaction with other protein(s).

Sporulation proteins appear to have mainly regulatory functions participating in protein-DNA, protein-protein and/or protein/PG interactions and only a small number of them have enzymatic activity. Many of these proteins could have lost their enzymatic activity and retained only substrate binding ability while some might have preserved the initial activity which is probably not directly relevant to their role in sporulation [72].

We were unable to purify BA1836 and perform biochemical characterization of this protein due to immediate proteolysis of the recombinant protein expressed in various *E. coli* strains (data not shown). This observation might indicate that interaction of BA1836 with other protein(s) is necessary for its stabilization against proteolysis.

The $\Delta ba1836$ mutant strain exhibited a reduced rate of germination which was partially complemented by *ba1836-gfp* expression (Figure 7). To determine whether this phenotype is due to the putative de-*N*-acetylase activity, we expressed a GFP-protein fusion version whereby the catalytic aspartic acid residue has been replaced with alanine, BA1836 (D53A) (Figure 7). Remarkably, the putative inactive BA1836 version partially complemented this phenotype, indicating that it is not caused by the lack of enzymatic activity. This result suggests that BA1836 may be an inactive deacetylase version which could participate in the organization of other enzymes or proteins involved in the germination process. Although BA1836 has the characteristic sequence motifs of active PDAs (Figure 1), we cannot exclude the possibility that mutations resulting in altering BA1836 conformation had occurred, and led to steric blocking of the active site or disruption of substrate-binding sites [73].

Absence of BA1836 did not change PG deacetylation either from stationary phase of growth or spores, suggesting that it is not a PGDA and therefore even if we managed to purify the protein, it is doubtful whether we would be able to demonstrate enzymatic activity on commonly used deacetylase substrates such as H³ glycol chitin, *N*-acetyl-chito-oligosaccharides and PG. This has been reported for PDAs acting on substrates other than PG, e.g. IcaB [74] and PgaB [75].

Therefore, BA1836 might have an as-yet-unidentified substrate, similarly to BA0330, and BA0331 (which also have the characteristic sequence motifs of active PDAs) [28], not directly relevant to its role in sporulation.

Presently studied putative and characterized PDAs from *B. anthracis* have been demonstrated to be involved in resistance to host lysozyme, stabilization of the cell wall, biogenesis of PG and for neutral polysaccharide modification and attachment to PG.

To our knowledge the sporulation specific role of BA1836 is unique among the functions of the other PDAs from *B. anthracis*.

Conflicts of Interest

The authors declare no conflicts of interest regarding the publication of this paper.

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Supporting Information

Table S1. List of strains and plasmids used in the present study.

Strains, plasmids	Description	Source/Reference
Strains		
<i>E. coli</i>		
DH5 α	<i>F</i> -80 <i>dlacZ</i> M15 (<i>lacZYA</i> - <i>argF</i>) <i>U169 recA1 endA1 hsdR17</i> (<i>rk</i> ⁻ , <i>mk</i> ⁺) <i>phoA</i> <i>supE44</i> - <i>thi</i> -1 <i>gyrA</i> 96 <i>relA</i> 1	Novagen
GM48	<i>thr</i> -1, <i>araC</i> 14, <i>leuB</i> 6 (Am), <i>fhuA</i> 31, <i>lacY</i> 1, <i>tsx</i> -78, <i>glnX</i> 44 (AS), <i>galK</i> 2 (Oc), <i>galT</i> 22, λ ⁻ , <i>dcm</i> -6, <i>dam</i> -3, <i>thiE</i> 1	Coli Genetic Stock Center (CGSC)
HB101	<i>F</i> - <i>hsd</i> -20 <i>recA</i> 13 <i>ara</i> -14 <i>proA</i> 2 <i>lacY</i> 1 <i>gal</i> K2 <i>rpsL</i> 20 (<i>Str</i>) <i>xyl</i> -5 <i>mtl</i> -1 <i>supE</i> 44	[36]
<i>B. anthracis</i>		
7702	pXO1 ⁺ , pXO2 ⁻	[35]
Plasmids		
pGEM T-Easy	Cloning vector	Promega
pNF8	pAT18 Ω (Pdl Ω gfp-mut1)	[38]
pWH1520	Cloning vector	Mobitec
pHT304-18Z	Promoterless <i>lacZ</i> vector	[41]
pRP1028	Tm ^S allelic-exchange vector	[37]
pSS4332	I-SceI expression vector	[37]

Table S2. List of oligonucleotides used in the present study.

Oligonucleotides	Sequence (5' \rightarrow 3')
<i>Peag</i> -FW	CCCAAGCTTTT TAGGTGATGAAGCGGAAAT
<i>Peag</i> -REV	CGGGATCCTTTATAAAATTCCTCCTTCAG
Up1836-pHT304-18Z-HindIII-FW	CCCAAGCTTCGTTATTTGCCCAAGATG
Up1836-pHT304-18Z-BamHI-REV	CGGGATCCAAAGCAACCTCACTATATTTT
<i>ba</i> 1836-FW-BamHI	CGGGATCCATGAAAAAGAAAATAATCATTACAATC
<i>ba</i> 1836-REV-NruI	TCGCGACGGGCCCGGGCCTTTTGCACTTTTTGTAAATTCATTTAC
<i>ba</i> 1836D53A-FW	TAACTTTTGCTGATGGCCC
<i>ba</i> 1836D53A-REV	GGGCCATCAGCAAAAGTTA
Upstream1836-FW	CCAAGCTTCCATATATCCAATACGGGCAAA
Upstream1836-REV	CCCCCGGGAACGATTGTAATGATTATTTCTTTTTC
Downstream1836-FW	CCCCCGGGAAGAAGGGCTATCAGTTCGTAACA
Downstream1836-REV	CGCGGATCCAGTTTACCGCGTTCAGCATT

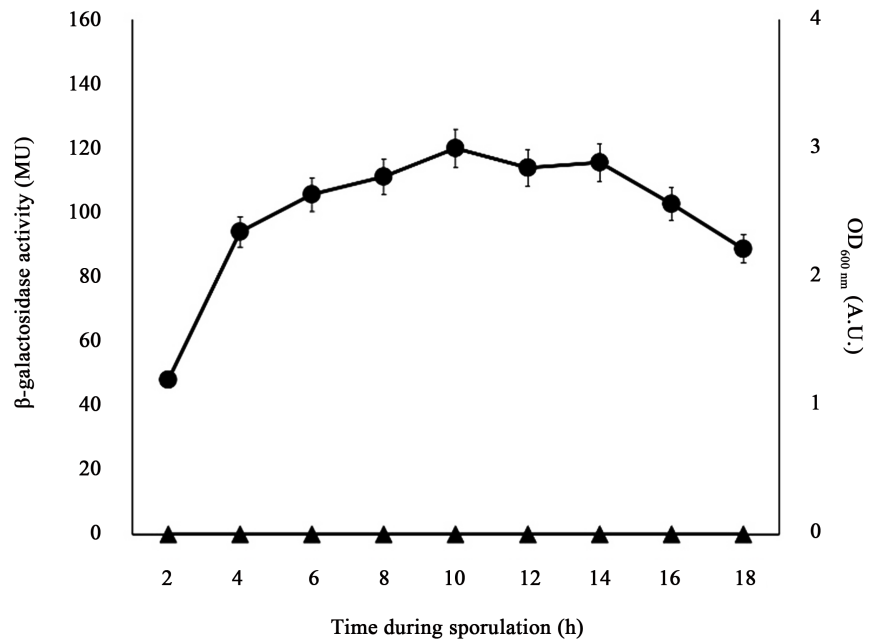


Figure S1. Expression of *ba1836* gene following induction of sporulation of exponentially phase cells into G modified medium. Samples taken at indicated time points were assayed for β -galactosidase activity. (●) $OD_{600\text{ nm}}$ (A.U.) and (▲) β -galactosidase activity of 7702 *B. anthracis* cells in Miller Units (MU).

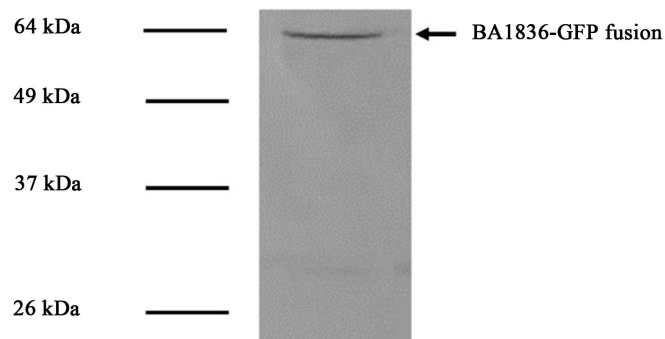


Figure S2. Western blot analysis of lysates of 7702 cells expressing *ba1836-gfp* during stationary phase of growth using a polyclonal anti-GFP antibody.

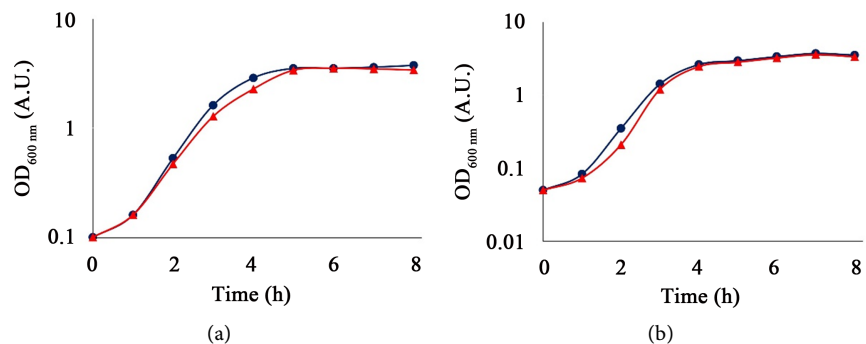


Figure S3. Growth curves of *B. anthracis* parental and mutant strains. Cultures of 7702 (●) and $\Delta ba1836$ mutant (▲) strains were grown in (a) BHI and (b) SPY liquid broth at 37°C. Cell growth was monitored at 600 nm.

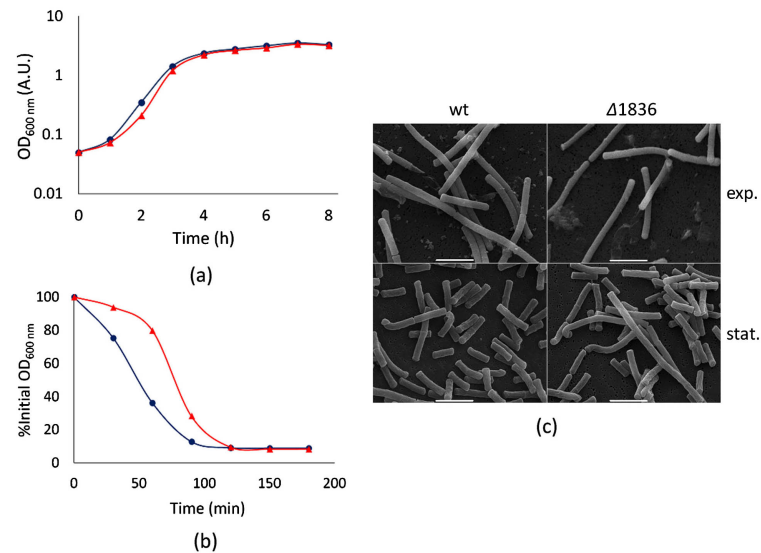


Figure S4. Lysozyme sensitivity, autolysis rate and morphology of parental and mutant strains. (a) Effect of lysozyme on 7702 (●) and $\Delta ba1836$ mutant (▲) strains. Strains were grown in SPY liquid broth at 37°C. $\Delta ba1836$ mutant is not affected by the addition of lysozyme; (b) Autolysis rate of 7702 (●) and $\Delta ba1836$ (▲) cells. Cell lysis was monitored by loss of absorbance at 600 nm. $\Delta ba1836$ strain is affected by autolysis. (c) SEM of wild type 7702 and $\Delta ba1836$ cells during exponential (exp.) and stationary (stat.) phase of growth. $\Delta ba1836$ cells exhibit longer chains of daughter cells compared to the wild type only during stationary phase of growth. Scale bar 5 μ m.

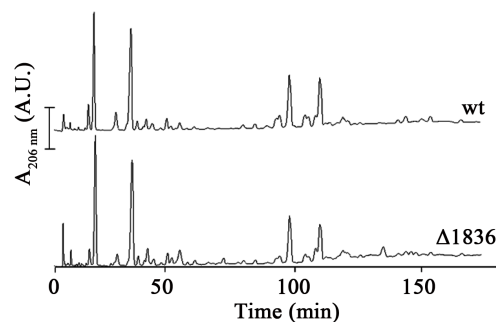


Figure S5. HPLC analysis of mucopeptide pattern of PG isolated from 7702 and $\Delta ba1836$ cells during stationary phase of growth. Bar, 200 mAU.

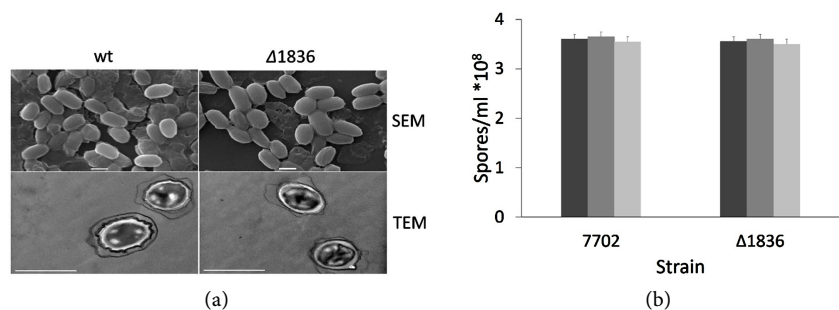


Figure S6. (a) Examination of morphology and structure of parental 7702 and mutant $\Delta ba1836$ mature spores using SEM and TEM. Scale bar 1 μ m. (b) Spore formation, viability and heat resistance of 7702 and $\Delta ba1836$ spores. (■) spores/ml (■) CFU/ml (■) CFU/ml after heat-treatment of spores.